

## Search Results -

	Terms						
***************************************	((IL-2 or interleukin-2) and (cancer or tumor or tumour)).clm. and (anergic or anergy)	16					



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## **Search History**

Today's Date: 4/13/2001

DB Name	Query	Hit Count	Set Name
USPT	((IL-2 or interleukin-2) and (cancer or tumor or tumour)).clm. and (anergic or anergy)	16	<u>L21</u>
USPT	((IL-2 or interleukin-2) and (cancer or tumor or tumour)).clm.	223	<u>L20</u>
USPT	(IL-2 or interleukin-2) and ((anergy or anergic) same (inhibit or inhibition or overcome or block or prevent or preventing or prevention or suppress\$))	161	<u>L19</u>
USPT	(IL-2 or interleukin-2) and (anergy or anergic)	303	<u>L18</u>
USPT	(IL-2 or interleukin-2).clm. and tumor and (anergy or anergic)	26	<u>L17</u>
USPT	(IL-2 or interleukin-2).clm. and tumor and anergy	25	<u>L16</u>
USPT	(IL-2 or interleukin-2) and tumor and anergy	223	<u>L15</u>
USPT	(IL-2 or interleukin-2) same (stimulate or stimulation or proliferate or proliferation or activate or activation) and IL-2.clm. and tumor.clm.	57	<u>L14</u>
USPT	(IL-2 or interleukin-2) same (stimulate or stimulation or proliferate or proliferation or activate or activation) and IL-2.clm. and tumor.clm	0	<u>L13</u>
USPT	(IL-2 or interleukin-2) same (stimulate or stimulation or proliferate or proliferation or activate or activation) and IL-2.clm. and anergy	18	<u>L12</u>
USPT	(IL-2 or interleukin-2) same (stimulate or stimulation or proliferate or proliferation or activate or activation) and IL-2.clm.	256	<u>L11</u>
USPT	(inhibit or inhibition or prevent or prevention or block) same anergy and method.clm.	74	<u>L10</u>
USPT	(inhibit or inhibition or prevent or prevention or block) same anergy	113	<u>L9</u>
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USPT	cytokine same gamma same anergy	10	<u>L7</u>
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USPT	BOUSSIOTIS-V-A.in.	0	<u>L5</u>
DWPI	BOUSSIOTIS-V-A.in.	4	<u>L4</u>
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DWPI	(modulating and gamma and cytokine).ti	18	<u>L2</u>
DWPI	Boussiotis-vassiliki-A.in.	0	<u>L1</u>







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		Tumor immun	ity as autoimmı	unity: tumor ant	igens includ	e normal self p	proteins which s
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			1993 Nov;14(11): PubMed - indexe]				
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	<b>114</b> .	Boussiotis VA. F	reeman G.J. Berezo	ovskaya A, Barber I	DL. Nadler LM		ı
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		Science. 1997 Oc	t 3;278(5335):124	l-8.	<i>5 5</i> -		
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	<b>5</b> :	Boussiotis VA, F	reeman GJ, Gribbe	n JG, Nadler LM.			F
		The role of B7	'-1/B7-2:CD28	CLTA-4 pathw	ays in the pi	revention of a	nergy, induction
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			96 Oct;153:5-26. I [PubMed - indexe]	Review. No abstracted for MEDLINE]	t available.		
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	<b>□6</b> :	Boussiotis VA, B Nadler LM.	arber DL, Nakarai	T, Freeman GJ, Gr	ibben JG, Berr	istein GM, D'And	lrea AD, Ritz J,
			T cell anergy by	y signaling throu	igh the gami	ma c chain of	the IL-2 recepto
		Science. 1994 No	ov 11;266(5187):10	039-42.			•
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	<b>7</b> :	Weissman IL.					F
		The common g		ι c) chain for mu			
			1994 Oct;85(10):	inside front cover. I	Review. No ab	stract available.	

PMID: 7961100 [PubMed - indexed for MEDLINE]

□8:	Grundstrom S, Dohlsten M, Sundstedt A.	R
	IL-2 unresponsiveness in anergic CD4+ T cells is due to defective signaling through the	е
	gamma-chain of the IL-2 receptor.	
	J Immunol. 2000 Feb 1;164(3):1175-84. PMID: 10640728 [PubMed - indexed for MEDLINE]	
	This. 10040720 [Ladvice medica for NEDSERVE]	
<b>9</b> :	Kaufman M, Andris F, Leo O. Free in PMC, F	?
	A logical analysis of T cell activation and anergy.	
	Proc Natl Acad Sci U S A. 1999 Mar 30;96(7):3894-9.	
	PMID: 10097134 [PubMed - indexed for MEDLINE]	
<b>10</b>	: Schwartz RH.	R
	Models of T cell anergy: is there a common molecular mechanism?	
	J Exp Med. 1996 Jul 1;184(1):1-8. Review. No abstract available.	
	PMID: 8691122 [PubMed - indexed for MEDLINE]	
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	Anergy as a mechanism of peripheral T cell tolerance.	
	J Immunol. 1996 Feb 15;156(4):1325-7. Review. No abstract available. PMID: 8568229 [PubMed - indexed for MEDLINE]	
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<b>12</b>	Selliah N, Finkel TH.	F
	Cutting edge: JAK3 activation and rescue of T cells from HIV gp120-induced unrespo	
	J Immunol. 1998 Jun 15;160(12):5697-701.	
	PMID: 9637477 [PubMed - indexed for MEDLINE]	
13	Boussiotis VA, Barber DL, Lee BJ, Gribben JG, Freeman GJ, Nadler LM.	F
	Differential association of protein tyrosine kinases with the T cell receptor is linked to	
	of anergy and its prevention by B7 family-mediated costimulation.	
	J Exp Med. 1996 Aug 1;184(2):365-76.	
	PMID: 8760790 [PubMed - indexed for MEDLINE]	
114	Boussiotis VA, Freeman GJ, Gribben JG, Nadler LM.	F
3 - T	The critical role of CD28 signalling in the prevention of human T-cell anergy.	
	Res Immunol. 1995 Mar-Apr;146(3):140-9. Review. No abstract available.	
	PMID: 8525043 [PubMed - indexed for MEDLINE]	
×****		_
15	fleeg K, Gaus II, Gifese D, Dendigs S, Whethice T, Wagner II.	F
	Superantigen-reactive T cells that display an anergic phenotype in vitro appear function Int Immunol. 1995 Jan;7(1):105-14.	
	PMID: 7718507 [PubMed - indexed for MEDLINE]	
<b>1</b> 6	Li XC, Demirci G, Ferrari-Lacraz S, Groves C, Coyle A, Malek TR, Strom TB.	F
	IL-15 and IL-2: a matter of life and death for T cells in vivo.	
	Nat Med. 2001 Jan;7(1):114-8.  PMID: 11135625 (PubMed_indexed for MEDLINE)	
	PMID: 11135625 [PubMed - indexed for MEDLINE]	
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Clear History

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13365401
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Interleukin-15 triggers the proliferation and cytotoxicity of granular lymphocytes in patients with lymphoproliferative disease of granular lymphocytes

Zambello R; Facco M; Trentin L; Sancetta R; Tassinari C; Perin A; Milani A; Pizzolo G; Rodeghiero F; Agostini C; Meazza R; Ferrini S; Semenzato G Univ. Padova, Dip. Med. Clin. Sperimentale, Via Giustiniani 2, 35128 Padova, Italy

Blood 89 (1). 1997. 201-211. Full Journal Title: Blood

ISSN: 0006-4971 Language: ENGLISH

Print Number: Biological Abstracts Vol. 103 Iss. 004 Ref. 053279 The recently cloned cytokine interleukin-15 (IL-15) shares several functional activities with IL-2 in different cell systems. Although IL-15 does not show sequence homology with IL-2, it uses components of the IL-2 receptor (IL-2R) for binding and signal transduction, namely, p75 (beta) and the p64 (gamma) chains of IL-2R. To evaluate whether IL-15 is involved of granular lymphocytes (GL) in patients with activation in lymphoproliferative disease of granular lymphocytes (LDGL), we evaluated the ability of IL-15 to stimulate GL proliferation, cytotoxic function, and the role of IL-2R beta and gamma molecules on relevant cells. Our results show that IL-15 stimulates cell proliferation and cytotoxic activity of GL in LDGL patients. Reverse-transcriptase polymerase chain reaction (RT-PCR) and phenotypic analyses using the anti-IL-2R gamma-chain -specific TUGh4 monoclonal antibody (MoAb) indicate that both CD3+ and CD3- GL express the p64 IL- $\overline{2}R$ , a result previously unknown. IL-15 activity was inhibited by antibodies against p75 and p64 IL-2R chains, while no inhibitory effects are detectable with anti-p55 IL-2R antibody. The association of anti-p75 and anti-p64 IL-2R MoAbs resulted in a nearly complete (95%) inhibition of IL-15-induced GL proliferation. Using RT-PCR analysis, we demonstrated that highly purified CD3+ and CD3- GL did not express mRNA for IL-15 or IL-2. By contrast, a clear-cut IL-15 mRNA signal was detected by RT-PCR in patients' peripheral blood mononuclear cells, with monocytes likely accounting for the source of IL-15 in LDGL patients. concentrated supernatants from enriched monocyte However, even in populations, we could not demonstrate the presence of IL-15 protein. Using anti-IL-15 specific MoAbs, a membrane-bound form of this cytokine was demonstrated both on CD3+ and CD3- LDGL cells. By RT-PCR analysis, purified  ${ t GL}$  from these patients were found to express the message for  ${ t IL}{ t -}15$  receptor a chain. Taken together, these results indicate that both CD3+ and CD3- GL are stimulated by IL-15 and that this cytokine mediates its activity through the beta and gamma chains of the IL-2R, providing further

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expansion in patients with LDGL.

13085365 BIOSIS Number: 99085365

IL-2 receptor gamma chain expression on CD34 positive hematopoietic progenitor cells from bone marrow and cord blood

Itano M; Tsuchiya S; Morita S; Fujie H; Ishii N; Yanagisawa T; Ohashi Y; Minegishi M; Sugamura K; Konno T

suggestions for the interpretation of the mechanisms that lead to cell

Dep. Pediatr. Oncol., Inst. Dev. Aging Cent., Tohoku Univ., Sendai 980-77, Japan

Tohoku Journal of Experimental Medicine 178 (4). 1996. 389-398. Full Journal Title: Tohoku Journal of Experimental Medicine

ISSN: 0040-8727 Language: ENGLISH

Print Number: Biological Abstracts Vol. 102 Iss. 004 Ref. 050814
The IL-2 receptor (IL-2R) gamma-chain is shared among receptors for IL-4,
IL-7, IL-9 and IL-15 as well as IL-2. In order to clarify the functional role of these cytokines interacting with the common ychain in human

early hematopoiesis, we studied expression of the IL-2R ychain on purified CD34 positive cells from bone marrow and cord blood. Broad populations of bone marrow mononuclear cells were all found to express the IL-2R gamma-chain . CD34 positive cells were purified by CD34 monoclonal antibodies and immunomagnetic beads as representative hematopoietic progenitor cells. It was established that only 38 +- 10% of CD34 positive bone marrow cells (n=5) and 35+-12% of CD34 positive cord blood cells (n=11) expressed the IL-2R ychain. CD34(+) IL-2R gamma-chain(+) and CD34(+) IL-2R gamma-chain(-) cells fractionated by cell sorting were subjected to clonogenic assays that showed granulocyte-macrophage colony-forming cells (CFU-GM) were present evenly in both fractions, whereas erythroid burst-forming cells (BFU-E) were enriched in the CD34(+) IL-2R gamma-chain(-) fraction approximately two- to six-fold as compared with CD34(+) IL-2R gamma-chian(+) fraction. Such clonogenic features did not differ between the bone marrow and cord blood cases. These results indicate that CD34(+) IL-2R gamma-chain(-) cells contain immature cells already committed to the erythroid lineage.

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13030859 BIOSIS Number: 99030859
Tolerance induction to human gamma

Tolerance induction to human gamma globulin in FcR gamma chain-deficient mice

Whitmer K J; Romball C G; Hobbs M V; Weigle W O Scripps Res. Inst., La Jolla, CA 92037, USA FASEB Journal 10 (6). 1996. A1179.

Full Journal Title: Joint Meeting of the American Society for Biochemistry and Molecular Biology, the American Society for Investigative Pathology and the American Association of Immunologists, New Orleans, Louisiana, USA, June 2-6, 1996. FASEB Journal

ISSN: 0892-6638 Language: ENGLISH

Print Number: Biological Abstracts/RRM Vol. 048 Iss. 007 Ref. 124595

3/7/4 (Item 4 from file: 55)
DIALOG(R)File 55:BIOSIS PREVIEWS(R)
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12130126 BIOSIS Number: 98730126

Transcript synthesis and surface expression of the interleukin-2 receptor (alpha-, beta-, and gamma-chain) by normal and malignant myeloid cells Schumann R R; Nakarai T; Gruss H-J; Brach M A; Von Arnim U; Kirschning C; Karawajew L; Ludwig W-D; Renauld J-C; Ritz J; Herrmann F

Humboldt Universitaet Berlin, Robert-Roessle Cancer Center, Lindenberger Weg 80, D-13122 Berlin, Germany

Blood 87 (6). 1996. 2419-2427.

Full Journal Title: Blood

ISSN: 0006-4971 Language: ENGLISH

Print Number: Biological Abstracts Vol. 101 Iss. 008 Ref. 114401

Expression of the interleukin-2 receptor alpha- (IL-2R-alpha), IL-2R-beta-, and the recently identified IL-2R-gamma-chain was examined on a wide range of cells of myeloid origin including neutrophils, monocytes, normal bone marrow-derived myeloid progenitors enriched for CD34+ cells, bone marrow blasts obtained from acute myelogenous leukemia (AML) patients, permanent myeloid leukemia cell lines by transcriptase-polymerase chain reaction and surface membrane analysis using receptor chain-specific monoclonal antibodies and flow cytometry. Expression of the p75 IL-2R-beta- and the p64 IL-2R-gamma-chain

was a common finding in most of the myeloid cell samples investigated, whereas IL-2R-alpha-chain was less frequently expressed. Although the

high-affinity IL-2R form (i.e., the alpha+, beta+, gamma+ IL-2R form) was detectable in a small minority of primary AML samples as well as the KG-1 cell line and IL-2 binding to these cells was sufficient to initiate signal transduction as evidenced by an increase in overall protein tyrosine phosphorylation and more specifically in tyrosine phosphorylation of the Janus kinase (JAK) 3, in none of these cell types did exposure to IL-2 affect cell growth kinetics. These results suggest that, in myeloid cells, the IL-2R may not stimulate mitogenic responses or that its components may be expressed in a combinational association with receptors for other cytokines and that IL-2R-gamma may play a regulatory role in normal and malignant myelopoiesis possibly independent from IL-2. Because recent studies by others have indicated that the IL-2R-gamma-chain may be shared by the IL-4R, the IL-7R, and most likely the IL-9R, expression of mRNA of these receptor types was also investigated in these cell samples. Surprisingly, in a substantial part of the myeloid lineage cells examined, an IL-2R-gamma+, IL-4R-, IL-7R- configuration was noted that was, however, frequently associated with expression of IL-9R. Sharing of IL-9R/IL-2R components was furthermore suggested by inhibition of 125I-IL-2 binding to primary AML cells with excess of unlabeled IL-9.

3/7/5 (Item 5 from file: 55)
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12096203 BIOSIS Number: 98696203

Reduced expression of the interleukin-2-receptor gamma chain on cord blood lymphocytes: Relationship to functional immaturity of the neonatal immune response

Zola H; Fusco M; Weedon H; Macardle P J; Ridings J; Roberton D M Child Health Res. Inst., Women's and Children's Hosp., 72 King William Rd., North Adelaide, SA 5006, Australia

Immunology 87 (1). 1996. 86-91. Full Journal Title: Immunology

ISSN: 0019-2805 Language: ENGLISH

Print Number: Biological Abstracts Vol. 101 Iss. 007 Ref. 096484 Mutation of the interleukin-2 (IL-2) receptor y chain, which also serves as a component of the receptor complexes for IL-4, 7, 9 and 15, results in severe immune deficiency. We hypothesized that the immunological immaturity of healthy neonates might be associated with low levels of expression of this receptor molecule. Using monoclonal antibody and a highly sensitive immunofluorescence method, we showed that IL-2 receptor gamma chain is expressed at significantly lower levels on cord blood cells compared with adult cells. IL-2-dependent T-cell activation in vitro was reduced in cord blood cells compared with adult cells, but B-cell responses to IL-4 were not obviously impaired. The lower level of expression of the y chain and some other cytokine receptor chains may contribute to the immunological immaturity of the newborn, by selectively depressing particular immunological mechanisms.

3/7/6 (Item 6 from file: 55)
DIALOG(R)File 55:BIOSIS PREVIEWS(R)
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11796675 BIOSIS Number: 98396675

Production of a mouse anti-human interleukin-2 receptor gamma chain specific monoclonal antibody

Holmberg P; Oetken C; Raivio E; Lindqvist C Abo Akademi Univ., Dep. Biochem, Abo, Finland 0 (0). 1995. 299.

Full Journal Title: 9TH INTERNATIONAL CONGRESS OF IMMUNOLOGY. The 9th International Congress of Immunology; Meeting Sponsored by the American Association of Immunologists and the International Union of Immunological

Societies, San Francisco, California, USA, July 23-29, 1995. ix+742p. 9th International Congress of Immunology: San Francisco, California, USA.

Language: ENGLISH

Print Number: Biological Abstracts/RRM Vol. 047 Iss. 009 Ref. 159078

3/7/7 (Item 7 from file: 55)
DIALOG(R)File 55:BIOSIS PREVIEWS(R)
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7000825 BIOSIS Number: 87061346

IDENTIFICATION OF THE INTERLEUKIN-2 RECEPTOR IL-2R ON HUMAN LEUKEMIC T CELLS USING COLLOIDAL GOLD AND SCANNING ELECTRON MICROSCOPY

HELINSKI E H; BIELAT K L; OVAK G M; MEENAGHAN M A; WIRTH J E; PAULY J L DEP. MOL. IMMUNOL., ROSWELL PARK MEML. INST., 666 ELM STREET, BUFFALO, N.Y. 14263.

J MED (WESTBURY) 19 (5-6). 1988. 353-368. CODEN: JNMDB Full Journal Title: Journal of Medicine (Westbury) Language: ENGLISH

Results of studies demonstrating the identification of the interleukin-2 receptor (IL-2R; i.e., anti-Tac) on the membrane ultra-structure of human leukemic T cells with an antibody carrying an electron dense colloidal gold microsphere (e.g., immiunogold) that was visualized using a scanning electron microscope (SEM) are reported. Our IL-2R model system employed HTLV-1 retrovirus-infected lymphoblastoid cells of the long-term human leukemic T cell line HUT-102B2. The presence of the IL-2R on these cells was defined using a double antibody procedure that employed as the primary atibody a purified mouse monoclonal anti-Leu-IL-2R antibody (mIgg1k, anti-Tac, CD25), and used as the secondary antibody a goat anti-mouse (gamma-chain specific) antibody that had been covalently bonded to a 40 nm colloidal gold particle. More than 95% of the  ${\tt HUT-102B2}$  were  ${\tt IL-2R+}$ , and there was a uniform distribution of the  ${\tt IL-2R}$ over the surface of the cells. Corresponding controls were employed in all examination and included IL-2R Jurkat human leukemic T cells and isotype identical immunoglobulins. The primary and secondary antibody reagents contained whole human serum and bovine serum albumin, and there was no evidence of the non-specific binding of these antibodies. These studies are the first to demonstrate the presence of a lymphokine receptor on the surface architecture of a cell. We anticipate no difficulty in applying the immunogold/SEM technology to define both normal and malignant cell membrane receptors for other cytokines.

3/7/8 (Item 1 from file: 72)
DIALOG(R)File 72:EMBASE
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9758068 EMBASE No: 95314770

Characterization of the IL-6 responsive elements in the gamma fibrinogen gene promoter

Zhang Z.; Fuentes N.L.; Fuller G.M.

Dept. of Cell Biology, University of Alabama, Birmingham, AL 35294-0005 USA

Journal of Biological Chemistry (USA) , 1995, 270/41 (24287-24291) CODEN: JBCHA  $\,$  ISSN: 0021-9258

LANGUAGES: English SUMMARY LANGUAGES: English

Fibrinogen, a hepatically derived class II acute phase protein, is the product of three separate genes, (Aalpha, Bbeta, and gamma). The fibrinogen genes are expressed constitutively; however, their transcription can he significantly up-regulated by interleukin-6 (IL-6) and glucocorticoid. Inspection of the promoter region of the fibrinogen gamma gene revealed three hexanucleotide clusters of CTGGGA that are recognized as class II IL-6 responsive elements. Functional analyses of these regions (designated here as site I, site II, and site III according to their position in the

promoter) were performed using luciferase reporter constructs and show a hierarchy of IL-6 response in which site II was the preferred functional site, site I was the next important site, and site III was the site least responsive to IL-6. Gel mobility shift assays using 25-base pair oligonucleotide probes derived from these three regions with the CTGGGA positioned in the middle and nuclear extracts from IL-6-treated primary complexes reveal the presence of IL-6-induced high molecular weight using anti-Stat3 antibody indicate that Stat3 is part of the IL-6-induced complex formed on the three gamma chain probes. The binding of Stat3 to the IL-6 responsive elements of the y probes is significantly weaker than to an alpha2-macroglobulin probe. These findings show for the first time that Stat3 is involved in associating with the IL-6 responsive elements of fibrinogen gamma chain, a class II acute phase gene other than alpha2-macroglobulin.

3/7/9 (Item 2 from file: 72)
DIALOG(R)File 72:EMBASE
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9398164 EMBASE No: 94336088

Expression of interleukin 2 receptors on human carcinoma cell lines and tumor growth inhibition by interleukin 2

Yasumura S.; Lin W.; Weidmann E.; Hebda P.; Whiteside T.L.

Pittsburgh Cancer Institute, W1041 Biomedical Science Tower, 211 Lothrop Street, Pittsburgh, PA 15213-2582 USA

INT. J. CANCER (USA) , 1994, 59/2 (225-234) CODEN: IJCNA ISSN:

LANGUAGES: English SUMMARY LANGUAGES: English

We have previously shown that human squamous cell carcinomas (SCC) express the interleukin 2 receptor (IL2R)-alpha and -beta chains, and that the ligand, IL2, directly inhibits growth of the tumor in vitro and in vivo in the tumor xenograft-nude mice model. We now show that the alpha and beta chains of IL2R are expressed on a variety of human carcinoma cell lines and on normal human keratinocytes in early-stage cultures. While all carcinoma in a population expressed IL2R-alpha and -beta proteins, keratinocytes obtained from different normal donors, variable proportions of cells were positive, as measured by flow cytometry. The carcinoma lines and 2/5 keratinocyte lines studied were also found to contain transcripts for the IL2R-beta chain detectable by combined reverse transcription-PCR (RTPCR) and hybridization with the specific cDNA probe. Incubation of the gastric (HR) or renal cell carcinoma (RCC) cell lines but not of other IL2R+ carcinoma cell lines or normal keratino- cytes, in the presence of IL2 resulted in dose-dependent inhibition of tumor cell growth. Monoclonal antibodies (MAbs) specific for IL2R-gamma chain completely reversed this growth inhibitory effect of IL2. The ligand, IL2, down-regulated surface expression of its own receptor and of intercellular adhesion molecule-I (ICAM-I) or class I major histocompatibility complex (MHC) antigens on IL2R+ tumor cells. All carcinoma cells studied incubated in the presence of IL2 exhibited significantly increased sensitivity to growth-inhibitory effects of other cytokines such as interferon (IFN)-gamma, tumor necrosis factor (TNF)-alpha or transforming growth factor (TGF)-beta IL2 inhibited growth of the HR cells by arresting a significant proportion of tumor cells in the GO/G1 phase of the cell cycle. Thus, IL2 can have direct effects on IL2R+ carcinoma cells, leading to changes in growth or to increases in sensitivity of tumor cells to cytostatic activities cytokines.

3/7/10 (Item 3 from file: 72)
DIALOG(R)File 72:EMBASE
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8760260 EMBASE No: 93063978

Principles of paramunization: Option and limits in veterinary medicine Buttner M.

Institute of Medical Microbiology, Infectious and Epidemic Diseases, Veterinary Faculty, Univ of Munich, Veterinarstrasse 13, 8000 Munchen 22 Germany

COMP. IMMUNOL. MICROBIOL. INFECT. DIS. (United Kingdom), 1993, 16/1 (1-10) CODEN: CIMID ISSN: 0147-9571

LANGUAGES: English SUMMARY LANGUAGES: English; French

The so-called primitive immune system has not changed during evolution. Even in primates it plays the most important role in first line defence against invading microorganisms. Cellular components such as macrophages, granulocytes, Natural Killer cells and gammadelta-T cells and soluble humoral factors - the **cytokines** - are the representants of the primitive immune system. An interlocking communicative network regulates flexible response of effector cells towards 'non-self' antigens. It also ensures close connection with the repertoire of specific immune response, e.g. antibody formation. Multifactorial diseases, nosocomial infections, diseases and forms of immunosuppression initiated various alternative methods in immunotherapy. Immunostimulation at the nonspecific defence level has first been noticed as 'side effects' of vaccination. Today it should be differentiated between substitution of the immune system with cytokines and induction of the non-specific defence repertoire mimicking natural antigen contact that is called paramunization. Advantages and disadvantages of both methods are discussed. In vitro as well as in vivo experiments with poxviruses document safety and efficacy of purified and inactivated virus particles in paramunization protocols. The main stimulative components of the poxvirus particles are located in the envelope of the virions. Poxvirus-induced stimulation of non-specific defence reactions is likely to have remote effects on the quality of further antigen processing. Besides the induction of a high short-term alertness in the primitive immune system paramunization may efficiently ongoing specific responses, e.g. immunoglobulin isotype influence selection. Therefore the term paramunization should not be used to characterize a separate part of the immune system, however, for didactic reasons it will facilitate the understanding of principles of the immune system.

3/7/11 (Item 1 from file: 154)
DIALOG(R)File 154:MEDLINE(R)
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06677496 90304313

Receptor expression and functional status of cultured human eosinophils derived from umbilical cord blood mononuclear cells.

Walsh GM; Hartnell A; Moqbel R; Cromwell O; Nagy L; Bradley B; Furitsu T; Ishizaka T; Kay AB

Department of Allergy and Clinical Immunology, National Heart and Lung Institute, London, UK.

Blood (UNITED STATES) Jul 1 1990, 76 (1) p105-11, ISSN 0006-4971 Journal Code: A8G

Contract/Grant No.: AI-10060, AI, NIAID

Languages: ENGLISH

Document type: JOURNAL ARTICLE

Selective use of recombinant human **cytokines** has enabled the culture of large numbers of eosinophils from human cord blood mononuclear cells, raising the possibility of their use as a model of eosinophil function. Cultured eosinophils (CE) were compared with normal-density peripheral blood eosinophils (PBE) in terms of their membrane receptor expression and function. Fc gamma R and CR1 expression of CE and PBE was similar. In contrast, the specific mean fluorescence for LFA-1 alpha, p150,95 alpha, ICAM-1, and HLA-DR was significantly elevated for CE compared with PBE. CE responded in PAF-induced chemotaxis in a similar fashion to PBE. CE gave higher numbers of both resting and platelet

activating factor (PAF)-stimulated immunoglobulin G (IqG) -C3b-dependent rosettes than PBE. CE and PBE had comparable capacity to kill IgG- and C-opsonized schistosomula in terms of both baseline values and PAF-induced enhancement of cytotoxicity. Baseline adherence by CE and PBE to plasma-coated glass was essentially the same, but stimulated adhesion (PAF) of CE was lower. Compared with PBE, CE generated less than half the amounts of extracellular and cell-associated PAF induced by calcium ionophore A23187 stimulation. Unlike PBE, CE did not generate PAF after exposure to IgG-coated Sepharose particles. CE stimulated with IgG-coated beads generated small quantities of LTC4, while A23187 stimulation resulted in approximately half the LTC4 levels observed with PBE. The total cell content of eosinophil peroxidase (EPO) was similar for CE and PBE. These data suggest that although CE and PBE have many phenotypic and functional properties in common there are quantitative differences that may be a consequence of their immaturity and/or the influence of the cytokines used in their culture.

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3/7/12
            (Item 1 from file: 399)
DIALOG(R) File 399:CA SEARCH(R)
(c) 1998 American Chemical Society. All rts. reserv.
  127016508
               CA: 127(2)16508y
                                  PATENT
  T cell proliferation and anergy regulation by interleukins or other
agents that stimulate cytokine receptor .gamma.-chains
  INVENTOR(AUTHOR): Boussiotis, Vassiliki A.; Nadler, Lee M.
  LOCATION: USA
  ASSIGNEE: Dana-Farber Cancer Institute
  PATENT: PCT International; WO 9717360 A2 DATE: 19970515
  APPLICATION: WO 96US17927 (19961112) *US 556038 (19951109)
  PAGES: 45 pp. CODEN: PIXXD2 LANGUAGE: English CLASS: C07K-000/A
  DESIGNATED COUNTRIES: AU; CA; JP DESIGNATED REGIONAL: AT; BE; CH; DE; DK
; ES; FI; FR; GB; GR; IE; IT; LU; MC; NL; PT; SE
  SECTION:
CA215005 Immunochemistry
CA201XXX Pharmacology
  IDENTIFIERS: T cell anergy regulation cytokine receptor, autoimmune
disease treatment cytokine receptor modulator, interleukin T cell anergy
  DESCRIPTORS:
Alloantigens... Antibodies... Antigens... Autoantigens... Tumor-associated
    agents that stimulating .gamma.-chain; t cell proliferation and anergy
    regulation by interleukins or other agents that stimulate cytokine
    receptor .gamma.-chains
Pathogenic bacteria...
    antigens that stimulating .gamma.-chain; t cell proliferation and
    anergy regulation by interleukins or other agents that stimulate
    cytokine receptor .gamma.-chains
Cytokine receptors... Interleukin 2 receptors... Interleukin 4 receptors...
Interleukin 7 receptors...
    .gamma.-chain regulation; t cell proliferation and anergy regulation by
    interleukins or other agents that stimulate cytokine receptor
    .gamma.-chains
Interleukin receptors...
   interleukin 15 receptors, .gamma.-chain regulation; t cell
   proliferation and anergy regulation by interleukins or other agents
   that stimulate cytokine receptor .gamma.-chains
Parasite... Virus...
   pathogenic, antigens that stimulating .gamma.-chain; t cell
   proliferation and anergy regulation by interleukins or other agents
   that stimulate cytokine receptor .gamma.-chains
   receptors, .gamma.-chain regulation; t cell proliferation and anergy
   regulation by interleukins or other agents that stimulate cytokine
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receptor .gamma.-chains
T-cell lymphopoiesis...
    regulation; t cell proliferation and anergy regulation by interleukins
    or other agents that stimulate cytokine receptor .gamma.-chains
Anergy... Drug screening... Immunotherapy... Interleukin 15... Interleukin
2... Interleukin 4... Interleukin 7...
    t cell proliferation and anergy regulation by interleukins or other
    agents that stimulate cytokine receptor .gamma.-chains
Autoimmune diseases... Graft-vs.-host reaction...
    treatment; t cell proliferation and anergy regulation by interleukins
    or other agents that stimulate cytokine receptor .gamma.-chains
  CAS REGISTRY NUMBERS:
157482-36-5 phosphorylation and assocn. with cytokine receptor
    .gamma.-chain; t cell proliferation and anergy regulation by
    interleukins or other agents that stimulate cytokine receptor
    .gamma.-chains
 3/7/13
            (Item 2 from file: 399)
DIALOG(R) File 399:CA SEARCH(R)
(c) 1998 American Chemical Society. All rts. reserv.
               CA: 125(7)84121a
                                   JOURNAL
  Single chain Ig/.gamma. gene-redirected human T lymphocytes produce
cytokines, specifically lyse tumor cells, and recycle lytic capacity
  AUTHOR(S): Weijtens, Mo E. M.; Willemsen, Ralph A.; Valerio, Dinko; Stam,
Kees; Bolhuis, Reinder L. H.
  LOCATION: Department Clinical Tumor Immunology, Daniel Hoed Cancer Center
, Rotterdam, Neth.
  JOURNAL: J. Immunol. DATE: 1996 VOLUME: 157 NUMBER: 2 PAGES: 836-843
  CODEN: JOIMA3 ISSN: 0022-1767 LANGUAGE: English
  SECTION:
CA215003 Immunochemistry
  IDENTIFIERS: tumor lysis T lymphocyte gene transduction, IgE Fc receptor
antibody transduction lymphocyte
  DESCRIPTORS:
Lymphokines and Cytokines, tumor necrosis factor-.alpha....
    cytokine formation by human T lymphocytes that have have a chimeric
    gene that encodes a single chain monoclonal antibody-IgE Fc receptor
    mol.
Immunoglobulin receptors, Fc. epsilon.RI (IgE fragment Fc receptor I)...
Receptors, Fc. epsilon.RI (IgE fragment Fc receptor I)...
    .gamma. chain; retrovirus-mediated transduction with a chimeric gene
    that encodes a single chain monoclonal antibody-IgE Fc receptor mol.
    stimulates human T lymphocytes to specifically lyse tumor cell
Therapeutics, geno-...
    retrovirus-mediated transduction of T-cells with a chimeric gene that
    encodes a single chain monoclonal antibody-IgE Fc receptor mol. in
    relation to
Kidney, neoplasm, renal cell carcinoma... Lymphocyte, T-cell, cytotoxic...
Neoplasm...
    retrovirus-mediated transduction with a chimeric gene that encodes a
    single chain Ig-IgE Fc receptor mol. stimulates human T lymphocytes to
    specifically lyse tumor cells
Antibodies, monoclonal... Cytolysis... Gene, animal, chimeric...
Transduction, genetic... Virus, animal, retro-...
    retrovirus-mediated transduction with a chimeric gene that encodes a
    single chain monoclonal antibody-IgE Fc receptor mol. stimulates human
    T lymphocytes to specifically lyse tumor cells
  CAS REGISTRY NUMBERS:
83869-56-1 cytokine formation by human T lymphocytes that have have a
```

chimeric gene that encodes a single chain monoclonal antibody-IgE Fc

receptor mol.

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(Item 3 from file: 399)
DIALOG(R) File 399:CA SEARCH(R)
(c) 1998 American Chemical Society. All rts. reserv.
               CA: 124(15)200210j
                                      PATENT
  Methods for modulating T cell responses by manipulating a common cytokine
receptor gamma chain
  INVENTOR (AUTHOR): Boussiotis, Vassiliki A.; Nadler, Lee M.
  LOCATION: USA
  ASSIGNEE: Dana-Farber Cancer Institute
  PATENT: PCT International; WO 9601122 A1 DATE: 960118
  APPLICATION: WO 95US8320 (950630) *US 270152 (940701)
  PAGES: 38 pp. CODEN: PIXXD2 LANGUAGE: English CLASS: A61K-038/20A;
A61K-039/395B; A61K-039/12B; A61K-039/02B; A61K-039/00B;
G01N-033/53B; G01N-033/68B; A61K-039/12J; A61K-038/20J; A61K-039/02K;
A61K-038/20K; A61K-039/002L; A61K-038/20L; A61K-039/00M; A61K-038/20M
  DESIGNATED COUNTRIES: AU; CA; JP DESIGNATED REGIONAL: AT; BE; CH; DE; DK
; ES; FR; GB; GR; IE; IT; LU; MC; NL; PT; SE
  SECTION:
CA215005 Immunochemistry
  IDENTIFIERS: T lymphocyte hematopoiesis cytokine receptor gamma,
interleukin 2 4 7 JAK kinase
  DESCRIPTORS:
Lymphokine and cytokine receptors... Receptors, cytokine...
    .gamma. chain; methods for modulating T cell responses by manipulating
    a common cytokine receptor gamma chain
Antigens... Antigens, allo-... Antigens, auto-... Antigens, tumor-assocd....
Autoimmune disease... Bacteria... Bone marrow, transplant...
Hematopoiesis, T-cell lymphopoiesis... Lymphokine and cytokine
receptors, interleukin 2... Lymphokine and cytokine receptors, interleukin 4
... Lymphokine and cytokine receptors, interleukin 7... Lymphokines and
Cytokines... Lymphokines and Cytokines, interleukin 2... Lymphokines and
Cytokines, interleukin 4... Lymphokines and Cytokines, interleukin 7...
Microorganism, pathogenic... Parasite... Receptors, interleukin 2...
Receptors, interleukin 4... Receptors, interleukin 7... Transplant and
Transplantation, allo-... Transplant and Transplantation, graft-vs.-host
reaction... Transplant and Transplantation, xeno-... Virus...
    methods for modulating T cell responses by manipulating a common
    cytokine receptor gamma chain
Antibodies...
    to cytokine receptor .gamma. chain; methods for modulating T cell
    responses by manipulating a common cytokine receptor gamma chain
  CAS REGISTRY NUMBERS:
161384-16-3 methods for modulating T cell responses by manipulating a
    common cytokine receptor gamma chain
 3/7/15
            (Item 4 from file: 399)
DIALOG(R) File 399:CA SEARCH(R)
(c) 1998 American Chemical Society. All rts. reserv.
  123110161
              CA: 123(9)110161u
                                    PATENT
  Antibody to human interleukin 2 receptor as immunosuppressant or
anti-allergic agent
  INVENTOR (AUTHOR): Shimamura, Toshiaki; Taki, Shinsuke; Hamuro, Junji;
Sugamura, Kazuo; Takeshita, Toshiichi; Kondo, Motonari
 LOCATION: Japan,
 ASSIGNEE: Ajinomoto Kk; Sugamura Kazuo
  PATENT: Japan Kokai Tokkyo Koho; JP 95149662 A2; JP 07149662 DATE:
950613
 APPLICATION: JP 94213706 (940907) *JP 93223574 (930908)
 PAGES: 11 pp. CODEN: JKXXAF LANGUAGE: Japanese CLASS: A61K-039/395A
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CA215003 Immunochemistry

IDENTIFIERS: interleukin 2 receptor gamma chain antibody,

```
immunosuppressant antiallergic monoclonal antibody IL2 receptor
   DESCRIPTORS:
 Allergy inhibitors... Antibodies, monoclonal...
 Glycophosphoproteins, interleukin 2-binding, p64... Immunosuppressants...
 Lymphokines and Cytokines, interleukin 4...
     monoclonal antibody to human interleukin 2 receptor .gamma. chain as
     immunosuppressant or anti-allergic agent that inhibits interleukin
     4-mediated disorder or allergy
  3/7/16
             (Item 5 from file: 399)
 DIALOG(R) File 399:CA SEARCH(R)
 (c) 1998 American Chemical Society. All rts. reserv.
   122158626
                CA: 122(13)158626f
                                      PATENT
  A monoclonal antibody to the interleukin 2 receptor .gamma. chain for use
 as an immunosuppressant
   INVENTOR (AUTHOR): Shimamura, Toshiro; Hamura, Junji; Nakazawa, Harumi;
 Kanayama, Yuka; Sugamura, Kazuo; Takeshita, Toshikazu
  LOCATION: Japan,
  ASSIGNEE: Ajinomoto Co., Inc.
  PATENT: European Pat. Appl. ; EP 621338 A2 DATE: 941026
  APPLICATION: EP 94106257 (940421) *JP 9394491 (930421) *JP 9436065
 (940307)
  PAGES: 37 pp. CODEN: EPXXDW LANGUAGE: English CLASS: C12N-015/13A;
C12P-021/08B; C12N-005/20B; A61K-039/395B; C12N-001/21B; C12N-005/10B;
C12P-021/00B DESIGNATED COUNTRIES: AT; BE; CH; DE; DK; ES; FR; GB; GR; IE;
IT; LI; LU; MC; NL; PT; SE
  SECTION:
CA215003 Immunochemistry
CA201XXX Pharmacology
  IDENTIFIERS: interleukin 2 receptor monoclonal antibody immunosuppressant
  DESCRIPTORS:
Allergy inhibitors... Antibodies, monoclonal... Immunosuppressants...
Inflammation inhibitors... Lymphokine and cytokine receptors, interleukin 2
... Receptors, interleukin 2...
    a monoclonal antibody to the human interleukin 2 receptor .gamma. chain
    for use as an immunosuppressant
Gene, animal...
    cDNA; a monoclonal antibody to the human interleukin 2 receptor .gamma.
    chain for use as an immunosuppressant
Deoxyribonucleic acid sequences, complementary...
    for fusion of proteins of light and heavy chains of monoclonal
    antibodies to human interleukin 2 receptor .gamma. subunit; a
    monoclonal antibody to the human interleukin 2 receptor .gamma. chain
    for u
Protein sequences...
    of fusion of proteins of light and heavy chains of monoclonal
    antibodies to human interleukin 2 receptor .gamma. subunit; a
    monoclonal antibody to the human interleukin 2 receptor .gamma. chain
    for us
Plasmid and Episome...
    pFv(GP-2)-DE, pFv(GP-4)-DE, cDNAs for Fv fragments of monoclonal
    antibody to .gamma. subunit of human interleukin 2 receptor; a
    monoclonal antibody to the human interleukin 2 receptor .gamma. chain
    fo
Plasmid and Episome...
   pIL2-RGS, cDNA for .gamma. subunit of human interleukin 2 receptor on,
    expression in Escherichia coli of; a monoclonal antibody to the human
    interleukin 2 receptor .gamma. chain for use as an immunosu
Autoimmune disease... Transplant and Transplantation, graft-vs.-host
reaction...
```

treatment of; a monoclonal antibody to the human interleukin 2 receptor .gamma. chain for use as an immunosuppressant CAS REGISTRY NUMBERS:

161309-68-8 161309-69-9 amino acid sequence; a monoclonal antibody to the human interleukin 2 receptor .gamma. chain for use as an

161309-70-2 161309-71-3 nucleotide sequence; a monoclonal antibody to the human interleukin 2 receptor .gamma. chain for use as an

3/7/17 (Item 6 from file: 399) DIALOG(R)File 399:CA SEARCH(R)

(c) 1998 American Chemical Society. All rts. reserv.

120268217 CA: 120(21)268217q PATENT

Cloning and expression of cDNA for human interleukin-2 receptor gamma

INVENTOR (AUTHOR): Sugamura, Kazuo; Takeshita, Toshikazu; Asao, Hironobu; Nakamura, Masataka; Shimamura, Toshiro; Suzuki, Manabu; Hamuro, Junji LOCATION: Japan.

ASSIGNEE: Ajinomoto Co., Inc.

PATENT: European Pat. Appl.; EP 578932 A2 DATE: 940119 APPLICATION: EP 93106561 (930422) \*JP 92104947 (920423)

PAGES: 50 pp. CODEN: EPXXDW LANGUAGE: English CLASS: C12N-015/12A; C07K-013/00B; C12P-021/08B; C12N-005/10B; G01N-033/48B; G01N-033/577B; A61K-037/02B; A61K-039/395B; A61K-039/44B DESIGNATED COUNTRIES: DE; FR; GB; IT

SECTION:

CA215005 Immunochemistry

CA203XXX Biochemical Genetics

IDENTIFIERS: interleukin 2 receptor gamma chain cDNA, cloning IL2 receptor gamma chain cDNA DESCRIPTORS:

Gene, animal...

cDNA, for interleukin-2 receptor gamma chain of human, cloning and expression of

Animal cell line, CHO... Animal cell line, L-929... Escherichia coli... Eukaryote... Prokaryote...

expression in, cDNA for human interleukin-2 receptor gamma chain for Deoxyribonucleic acid sequences, complementary...

for interleukin-2 receptor gamma chain of human

Lymphokines and Cytokines, interleukin 2, receptors... Receptors, interleukin 2...

gamma chain of, cDNA for, of human, cloning and expression of  ${\tt Immunomodulators...}$ 

human interleukin-2 receptor gamma chain and its antibody as Protein sequences...

of interleukin-2 receptor gamma chain of human

Antibodies... Antibodies, monoclonal...

to human interleukin-2 receptor gamma chain, prepn. of CAS REGISTRY NUMBERS:

148348-31-6 154609-72-0 amino acid sequence of and cloning and expression of cDNA for

154609-71-9 154609-73-1 154609-74-2 154609-75-3 154609-76-4 nucleotide sequence and cloning of

## ? e au=boussiotis

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Items Index-term
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16 AU=BOUSSIOS, THALIA
  E2
  E3
           0 *AU=BOUSSIOTIS
 E4
           11 AU=BOUSSIOTIS V
 E5
           71 AU=BOUSSIOTIS V A
 E6
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 E7
          44 AU=BOUSSIOTIS V.A.
39 AU=BOUSSIOTIS VA
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2 AU=BOUSSIOTIS, VASSILIKI
23 AU=BOUSSIOTIS, VASSILIKI A.
2 AU=BOUSSIOTIS, VICKI A.
 E9
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 E12
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       5 AU=BOUSSIOTIS, VIKI A.
 E14
           1 AU=BOUSSIOTOY A
 E15
          10 AU=BOUSSIOU M
 E16
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 E17
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E18
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E19
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E21
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E23
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E24
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               71 AU=BOUSSIOTIS V A
                1 AU=BOUSSIOTIS V.
               44 AU=BOUSSIOTIS V.A.
               39 AU=BOUSSIOTIS VA
                1 AU=BOUSSIOTIS, V. A.
                2 AU=BOUSSIOTIS, VASSILIKI
               23 AU=BOUSSIOTIS, VASSILIKI A.
              191 E4-E11
? s s4 and (gamma(w)chain
>>>Unmatched parentheses
? s s4 and gamma(w)chain
              191 S4
          516643 GAMMA
          678765 CHAIN
6405 GAMMA(W)CHAIN
18 S4 AND GAMMA(W)CHAIN
      S5
? rd s5
```

13090260 BIOSIS Number: 99090260

Pre-B acute lymphoblastic leukemia cells may induce T-cell anergy to alloantigen

Cardoso A A; Schultze J L; Boussiotis V A; Freeman G J; Seamon M J; Laszlo S; Billet A; Sallan S E; Gribben J G; Nadler L M

Dana-Farber Cancer Inst., D-740, 44 Binney St. Boston, MA 02115, USA Blood 88 (1). 1996. 41-48.

Full Journal Title: Blood

ISSN: 0006-4971 Language: ENGLISH

Print Number: Biological Abstracts Vol. 102 Iss. 004 Ref. 055709 Even if neoplastic cells express tumor associated antigens they still may fail to function as antigen presenting cells (APC) if they lack expression of one or more molecules critical for the induction of productive immunity. defects can be repaired by physiologic activation, cellular transfection, or fusion of tumor cells with professional APC. Although such defects can be repaired, antitumor specific T cells may still fail to respond in vivo if they may have been tolerized. Here, human pre-B cell acute lymphoblastic leukemia (pre-B ALL) was used as a model to determine if primary human tumor cells can function as alloantigen presenting cells (alloAPC) or alternatively whether they induce energy. In the present report, we show that pre-B cell ALL express alloantigen and adhesion molecules but uniformly lack B7-1 (CD80) and only a subset express B7-2 (CD86). Pre-B ALL cells are inefficient or ineffective alloAPC and those cases that lack expression of B7-1 and B7-2 also induce alloantigen unresponsiveness. Under these circumstances, T-cell T-cell unresponsiveness could be prevented by physiologic activation of tumor cells via CD40, cross-linking CD28, or signaling through the common gamma chain of the interleukin-2 receptor on T cells. Taken together, these results suggest that pre-B ALL may be incapable of inducing clinically significant T-cell-mediated antileukemia responses. This defect may be not only due to their inability to function as APC, but also due to potential to induce tolerance. Attempts to induce clinically significant antitumor immune responses may then require not only mechanisms to repair the antigen presenting capacity of the tumor cells, but also reversal of tolerance.

6/7/2 (Item 2 from file: 55)
DIALOG(R)File 55:BIOSIS PREVIEWS(R)
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13039673 BIOSIS Number: 99039673

Complete blockade of B7 family-mediated costimulation is necessary to induce human alloantigen-specific anergy: A method to ameliorate graft-versus-host disease and extend the donor pool

Gribben J G; Guinan E C; Boussiotis V A; Ke X-Y; Linsle L; Sieff C; Gray G S; Freeman G J; Nadler L M

Div. Hematol. Malignancies, Dana-Farber Cancer Inst., 44 Binney St., Boston, MA 02115, USA

Blood 87 (11). 1996. 4887-4893.

Full Journal Title: Blood

ISSN: 0006-4971

Language: ENGLISH Print Number: Biological Abstracts Vol. 102 Iss. 002 Ref. 021846

Graft-versus-host disease (GVHD) is initiated by adoptively transferred donor T cells that recognize host alloantigens. Whereas the absence of donor T-cell proliferation to host alloantigens in a mixed-leukocyte reaction does not predict freedom from GVHD, the frequency of alloreactive precursor helper T lymphocytes (pHTL) is predictive. Complete blockade of B7 family-mediated costimulation, but not of major histocompatibility complex recognition or adhesion, induces host alloantigen-specific energy by reducing cytokine production below threshold levels necessary for common gamma-chain signaling. The associated reduction of alloreactive pHTL frequency below that predictive for GVHD, without depletion of either nonallospecific T cells or hematopoietic progenitors, has led us to embark upon human clinical trials of haplomismatched allogeneic bone marrow transplantation.

6/7/3 (Item 3 from file: 55) DIALOG(R) File 55:BIOSIS PREVIEWS(R) (c) 1998 BIOSIS. All rts. reserv.

11941641 BIOSIS Number: 98541641

The critical role of CD28 signalling in the prevention of human T-cell

Boussiotis V A; Freeman G J; Gribben J G; Nadler L M

Div. Hematol. Malignancies, Dana Farber Cancer Inst., Harv. Med. Sch., Boston, MA 02115, USA

Research in Immunology 146 (3). 1995. 140-149.

Full Journal Title: Research in Immunology

ISSN: 0923-2494 Language: ENGLISH

Print Number: Biological Abstracts Vol. 100 Iss. 012 Ref. 178332

6/7/4 (Item 4 from file: 55) DIALOG(R) File 55:BIOSIS PREVIEWS(R) (c) 1998 BIOSIS. All rts. reserv.

11469081 BIOSIS Number: 98069081

Common gamma-chain signaling is sufficient to prevent

alloantigen specific T-cell clonal anergy

Boussiotis V A; Barber D L; Nakarai T; Freeman G J; Gribben J G; Bernstein G M; D'Andrea A D; Ritz J; Nadler L M

Div. Hematologic Malignancies Pediatric Oncol., Dana-Farber Cancer Inst., Harvard Med. Sch., Boston, MA, USA

Blood 84 (10 SUPPL. 1). 1994. 111A.

Full Journal Title: Abstracts Submitted to the 36th Annual Meeting of the American Society of Hematology, Nashville, Tennessee, USA, December 2-6, 1994. Blood

ISSN: 0006-4971 Language: ENGLISH

Print Number: Biological Abstracts/RRM Vol. 047 Iss. 002 Ref. 030668

6/7/5 (Item 5 from file: 55) DIALOG(R) File 55:BIOSIS PREVIEWS(R) (c) 1998 BIOSIS. All rts. reserv.

11435459 BIOSIS Number: 98035459

Prevention of T cell anergy by signaling through the gamma-c chain of the

Boussiotis V A; Barber D L; Nakarai T; Freeman G J; Gribben J G; Bernstein G M; D'Andrea A D; Ritz J; Nadler L M

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Science (Washington D C) 266 (5187). 1994. 1039-1042. Full Journal Title: Science (Washington D C)

ISSN: 0036-8075 Language: ENGLISH

Print Number: Biological Abstracts Vol. 099 Iss. 002 Ref. 020003 stimulated through their antigen receptor without requisite costimulation, T cells enter a state of antigen-specific unresponsiveness, termed anergy. In this study, signaling through the common gamma chain of the interleukin-2 (IL-2), IL-4, and IL-7 receptors in the presence of antigen was found to be sufficient to prevent the induction of anergy. After culture with IL-2, IL-4, or IL-7, Jak3 kinase was tyrosine-phosphorylated, which correlated with the prevention of anergy. Therefore, a signal through the common gamma chain may regulate the decision of T cells to either clonally expand or enter a state of

6/7/6 (Item 1 from file: 72) DIALOG(R)File 72:EMBASE (c) 1998 Elsevier Science B.V. All rts. reserv.

9616444 EMBASE No: 95171678

B7-1 and B7-2 do not deliver identical costimulatory signals, since B7-2but not B7-1 preferentially costimulates the initial production of IL-4Freeman G.J.; Boussiotis V.A.; Anumanthan A.; Bernstein G.M.; Ke

X.-Y.; Rennert P.D.; Gray G.S.; Gribben J.G.; Nadler L.M.

Dana-Farber Cancer Institute, Department of Medicine, Harvard Medical School, Boston, MA 02115 USA

Immunity (USA) , 1995, 2/5 (523-532) CODEN: IUNIE ISSN: 1074-7613 LANGUAGES: English SUMMARY LANGUAGES: English

The functional necessity for two CD28 counterreceptors (B7-1 and B7-2) is presently unknown. B7-1 and B7-2 equivalently costimulate IL-2 and interferon-gamma (IFNgamma) production and IL-2 receptor alpha and gamma chain expression. B7-2 induces significantly more IL-4 production than B7-1, with the greatest difference seen in naive T cells. Repetitive costimulation of CD4+CD45RA+ T cells with B7-2 results in moderate levels of both IL-4 and IL-2, whereas repetitive costimulation with B7-1 results in high levels of IL-2 and low levels of IL-4. Therefore, B7-1 and B7-2 costimulation mediate distinct outcomes, since B7-2 provides an initial signal to induce naive T cells to become IL-4 producers, thereby directing the immune response more towards ThO/Th2, whereas B7-1 is a more neutral differentiative signal.

6/7/7 (Item 1 from file: 399) DIALOG(R) File 399:CA SEARCH(R) (c) 1998 American Chemical Society. All rts. reserv.

127016508 CA: 127(2)16508y PATENT

T cell proliferation and anergy regulation by interleukins or other agents that stimulate cytokine receptor .gamma.-chains

INVENTOR (AUTHOR): Boussiotis, Vassiliki A.; Nadler, Lee M.

LOCATION: USA

ASSIGNEE: Dana-Farber Cancer Institute

PATENT: PCT International; WO 9717360 A2 DATE: 19970515 APPLICATION: WO 96US17927 (19961112) \*US 556038 (19951109)

PAGES: 45 pp. CODEN: PIXXD2 LANGUAGE: English CLASS: C07K-000/A DESIGNATED COUNTRIES: AU; CA; JP DESIGNATED REGIONAL: AT; BE; CH; DE; DK

; ES; FI; FR; GB; GR; IE; IT; LU; MC; NL; PT; SE SECTION:

CA215005 Immunochemistry CA201XXX Pharmacology

IDENTIFIERS: T cell anergy regulation cytokine receptor, autoimmune disease treatment cytokine receptor modulator, interleukin T cell anergy

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DESCRIPTORS:
  Alloantigens... Antibodies... Antigens... Autoantigens... Tumor-associated
      agents that stimulating .gamma.-chain; t cell proliferation and anergy
      regulation by interleukins or other agents that stimulate cytokine
      receptor .gamma.-chains
  Pathogenic bacteria...
      antigens that stimulating .gamma.-chain; t cell proliferation and
      anergy regulation by interleukins or other agents that stimulate
      cytokine receptor .gamma.-chains
  Cytokine receptors... Interleukin 2 receptors... Interleukin 4 receptors...
  Interleukin 7 receptors...
      .gamma.-chain regulation; t cell proliferation and anergy regulation by
      interleukins or other agents that stimulate cytokine receptor
      .gamma.-chains
  Interleukin receptors...
     interleukin 15 receptors, .gamma.-chain regulation; t cell
     proliferation and anergy regulation by interleukins or other agents
     that stimulate cytokine receptor .gamma.-chains
 Parasite... Virus...
     pathogenic, antigens that stimulating .gamma.-chain; t cell
     proliferation and anergy regulation by interleukins or other agents
     that stimulate cytokine receptor .gamma.-chains
 Interleukin 15...
     receptors, .gamma.-chain regulation; t cell proliferation and anergy
     regulation by interleukins or other agents that stimulate cytokine
     receptor .gamma.-chains
 T-cell lymphopoiesis...
     regulation; t cell proliferation and anergy regulation by interleukins
     or other agents that stimulate cytokine receptor .gamma.-chains
 Anergy... Drug screening... Immunotherapy... Interleukin 15... Interleukin
 2... Interleukin 4... Interleukin 7...
     t cell proliferation and anergy regulation by interleukins or other
     agents that stimulate cytokine receptor .gamma.-chains
 Autoimmune diseases... Graft-vs.-host reaction...
     treatment; t cell proliferation and anergy regulation by interleukins
     or other agents that stimulate cytokine receptor .gamma.-chains
   CAS REGISTRY NUMBERS:
 157482-36-5 phosphorylation and assocn. with cytokine receptor
     .gamma.-chain; t cell proliferation and anergy regulation by
    interleukins or other agents that stimulate cytokine receptor
     .gamma.-chains
 6/7/8
            (Item 2 from file: 399)
DIALOG(R) File 399:CA SEARCH(R)
(c) 1998 American Chemical Society. All rts. reserv.
  124200210
               CA: 124(15)200210j
                                     PATENT
  Methods for modulating T cell responses by manipulating a common cytokine
receptor gamma chain
  INVENTOR (AUTHOR): Boussiotis, Vassiliki A.; Nadler, Lee M.
  LOCATION: USA
  ASSIGNEE: Dana-Farber Cancer Institute
  PATENT: PCT International ; WO 9601122 A1 DATE: 960118
  APPLICATION: WO 95US8320 (950630) *US 270152 (940701)
  PAGES: 38 pp. CODEN: PIXXD2 LANGUAGE: English CLASS: A61K-038/20A;
A61K-039/395B; A61K-039/12B; A61K-039/02B; A61K-039/00B;
G01N-033/53B; G01N-033/68B; A61K-039/12J; A61K-038/20J; A61K-039/02K;
A61K-038/20K; A61K-039/002L; A61K-038/20L; A61K-039/00M; A61K-038/20M
  DESIGNATED COUNTRIES: AU; CA; JP DESIGNATED REGIONAL: AT; BE; CH; DE; DK
; ES; FR; GB; GR; IE; IT; LU; MC; NL; PT; SE
  SECTION:
CA215005 Immunochemistry
 IDENTIFIERS: T lymphocyte hematopoiesis cytokine receptor gamma,
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interleukin 2 4 7 JAK kinase DESCRIPTORS: Lymphokine and cytokine receptors... Receptors, cytokine... .gamma. chain; methods for modulating T cell responses by manipulating a common cytokine receptor gamma chain Antigens... Antigens, allo-... Antigens, auto-... Antigens, tumor-assocd.... Autoimmune disease... Bacteria... Bone marrow, transplant... Hematopoiesis, T-cell lymphopoiesis... Lymphokine and cytokine receptors, interleukin 2... Lymphokine and cytokine receptors, interleukin 4 ... Lymphokine and cytokine receptors, interleukin 7... Lymphokines and Cytokines... Lymphokines and Cytokines, interleukin 2... Lymphokines and Cytokines, interleukin 4... Lymphokines and Cytokines, interleukin 7... Microorganism, pathogenic... Parasite... Receptors, interleukin 2... Receptors, interleukin 4... Receptors, interleukin 7... Transplant and Transplantation, allo-... Transplant and Transplantation, graft-vs.-host reaction... Transplant and Transplantation, xeno-... Virus... methods for modulating T cell responses by manipulating a common cytokine receptor gamma chain Antibodies... to cytokine receptor .gamma. chain; methods for modulating T cell responses by manipulating a common cytokine receptor gamma chain CAS REGISTRY NUMBERS: 161384-16-3 methods for modulating T cell responses by manipulating a common cytokine receptor gamma chain 6/7/9 (Item 1 from file: 351) DIALOG(R) File 351: DERWENT WPI (c)1998 Derwent Info Ltd. All rts. reserv. 011303072 \*\*Image available\*\* WPI Acc No: 97-280977/199725 Stimulating or inhibiting proliferation of T cells expressing cytokine receptor gamma chain - comprises treatment with e.g. antibody that binds to this chain, useful for treatment of e.g. auto-immune disease, transplant rejection and guest versus host diseases Patent Assignee: DANA FARBER CANCER INST INC (DAND ) Inventor: BOUSSIOTIS V A; NADLER L M Number of Countries: 020 Number of Patents: 002 Patent Family: Patent No Kind Date Applicat No Kind Date Main IPC WO 9717360 A2 19970515 WO 96US17927 A 19961112 Week 199725 B AU 9710506 A 19970529 AU 9710506 A 19961112 199737 Priority Applications (No Type Date): US 95556038 A 19951109 Cited Patents: No-SR.Pub Patent Details: Patent Kind Lan Pg Filing Notes Application Patent WO 9717360 A2 E 46 Designated States (National): AU CA JP Designated States (Regional): AT BE CH DE DK ES FI FR GB GR IE IT LU MC NL PT SE AU 9710506 A Based on WO 9717360 Abstract (Basic): WO 9717360 A Stimulating proliferation of T cells that express a cytokine receptor gamma chain (A), and which have received a primary activating signal under conditions that normally result in unresponsiveness comprises treating the T cells with an agent (I) that binds (A) and stimulates an intracellular signal in the cell that causes proliferation. (I) is not natural interleukin (IL)-2. Also new

are: (1) induction of unresponsiveness to an antigen (Ag) in a T cell that expresses (A) by contacting it, in presence of Ag, with an agent (II) that inhibits delivery of a signal through (A); (2) a method for

identifying (II).

USE - Induction of unresponsiveness is useful for treating a wide variety of autoimmune diseases, transplant rejection, unwanted immune responses such as allergies and especially graft versus host disease in patients given bone marrow transplants. Inducing proliferation is also used to improve the response to a vaccinating Ag, derived from a microorganism or tumour.

Dwg.1/11

Derwent Class: B04

International Patent Class (Main): C07K-000/00; C07K-014/00

6/7/10 (Item 2 from file: 351)

DIALOG(R) File 351: DERWENT WPI

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010590567 \*\*Image available\*\*

WPI Acc No: 96-087520/199609

Modulation of T cell responses with therapeutic applications - by

manipulating a common cytokine receptor gamma chain

Patent Assignee: DANA FARBER CANCER INST INC (DAND )

Inventor: BOUSSIOTIS V A; NADLER L M

Number of Countries: 020 Number of Patents: 003

Patent Family:

Patent No Kind Date Applicat No Kind Date WO 9601122 A1 19960118 WO 95US8320 A 19950630 A61K-038/20 Week AU 9529152 A 19960125 AU 9529152 A 19950630 A61K-038/20 EP 768890 A1 19970423 EP 95924766 A 19950630 A61K-038/20 WO 95US8320 A 19950630 199609 B AU 9529152 A 19960125 AU 9529152 199618 199721

Priority Applications (No Type Date): US 94270152 A 19940701

Cited Patents: 06Jnl.Ref; EP 621338

Patent Details:

Patent Kind Lan Pg Filing Notes Application Patent

WO 9601122 A1 E 39

Designated States (National): AU CA JP

Designated States (Regional): AT BE CH DE DK ES FR GB GR IE IT LU MC NL PT SE

AU 9529152 A Based on

WO 9601122 EP 768890 A1 E Based on WO 9601122

Designated States (Regional): AT BE CH DE DK ES FR GB GR IE IT LI LU MC NL PT SE

Abstract (Basic): WO 9601122 A

The following are claimed: (A) a method for stimulating proliferation by a T cell which expresses a cytokine receptor gamma-chain and which has received a prim. activation signal under conditions, which normally result in no response in a T cell, comprises contacting the T cell with an agent (I) which binds to the cytokine receptor gamma-chain and stimulates an intracellular signal in the T cell resulting in T cell proliferation, provided that (I) does not consist of natural interleukin (I1)-2; (B) a method in which (I) acts intracellularly to stimulate phosphorylation of a JAK kinase with a mol.wt. of about 116kD as determined by SDS polyacrylamide gel electrophoresis, resulting in proliferation of the T cell; (C) a method for inducing no response to an antigen (Ag) in a T cell which expresses a cytokine receptor gamma-chain, comprising contacting the T cell in the presence of an Ag with an agent (II), which inhibits delivery of a signal through the cytokine receptor gamma-chain, resulting in T cell no response to the antigen; (D) a method for inhibiting graft-versus-host disease (GVHD) in a bone marrow transplant recipient by contacting a donor T cell, which expresses a cytokine receptor gamma-chain, with a cell which expresses a recipient Ag and (II), resulting in donor T cell no response to the cell which expresses the recipient Ag; and (E) a method for identifying (II) comprising (a) contacting a T cell which expresses

a cytokine receptor <code>gamma-chain</code> with (1) a first agent which stimulates a prim. activation signal in the T cell, (2) a second agent which stimulates an intracellular signal through the cytokine receptor <code>gamma-chain</code> and (3) a third agent to be tested for the ability to inhibit delivery of the signal through the cytokine receptor <code>gamma-chain</code>, and (b) determining the presence of T cell proliferation, in which inhibition of T cell proliferation indicates that the third agent inhibits delivery of a signal to T cell through the cytokine receptor <code>gamma-chain</code>.

USE - (I) may be used to enhance an anti-tumour response or a T-cell response to pathogens, such as viruses, bacteria, fungi and parasites. (I) may also be used to increase the efficacy of vaccination. (II) may be used to prevent organ transplant rejection and to inhibit GVHD, to treat autoimmune diseases such as diabetes mellitus, arthritis, multiple sclerosis, SLE, dermatitis, psoriasis, Sjogren's Syndrome, alopecia areata, Crohn's disease, asthma and vaginitis, to treat allergy and allergic reactions and to energise T Dwg.1/4

Derwent Class: B04; D16; S03

International Patent Class (Main): A61K-038/20

International Patent Class (Additional): A61K-039/00; A61K-039/002; A61K-039/12; A61K-039/395; G01N-033/53; G01N-033/68;